

GenCore version 5.1.3
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OM nucleic - nucleic search, using sw model

Run on: October 23, 2002, 19:00:33 ; Search time 179 Seconds
(without alignments)
3347.504 Million cell updates/sec

Title: US-09-728-446-819
Perfect score: 349
Sequence: 1 tattatatgtaagtaacnctg.....gnntggccttgaaggttg 349

Scoring table: IDENTITY_NUC
Gapop 10.0 , Gapext 1.0

Searched: 1736436 seqs, 858457221 residues
Total number of hits satisfying chosen parameters: 3472872

Minimum DB seq length: 0
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

Database : N_Geneseq_032802:*

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14:	/SIDS1/gcgdata/geneseq/geneseqn-emb1/NA1993.DAT:*
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22:	/SIDS1/gcgdata/geneseq/geneseqn-emb1/NA2001A.DAT:*
23:	/SIDS1/gcgdata/geneseq/geneseqn-emb1/NA2001B.DAT:*
24:	/SIDS1/gcgdata/geneseq/geneseqn-emb1/NA2002.DAT:*

Pred. NO. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	130.6	37.4	1209	19	AAV32579 Human high voltage
2	130.6	37.4	1393	18	AAV94469 Human Fchd545 gene
3	130.6	37.4	1393	21	AAZ50709 Nucleotide sequenc
4	130.6	37.4	1414	20	AAV57719 Voltage-dependent
5	105	34.7	1062	23	AAV74839 DNA encoding novel
6	105	30.1	1672	23	AAV74840 DNA encoding novel
7	77.8	22.3	435	21	AAC01263 Human secreted pro
8	57.8	16.6	473	23	AAV74836 DNA encoding novel
9	51	14.6	2698	17	AAV08126 Mouse synecan-1 g

10	51	14.6	26700	15	AAO67902	Syndecan gene. Mu
11	51	14.6	26700	19	AAV15946	Mouse syndecan gen
12	51	14.6	26700	20	AAV81283	Mouse syndecan-1 e
13	50	14.3	178	20	AAV85695	Novel CDNA sequenc
14	50	14.3	5889	20	AAV84328	Mouse A-myb genomi
15	50	14.3	6094	22	AAV63435	Murine CD39-L2 gen
16	49.4	14.2	14775	24	AB199535	Mouse ischaemic co
17	49.4	14.2	36901	20	AAZ23897	Murine LOBO genomi
18	49.4	14.2	38886	20	AAZ23897	Murine LOBO genomi
19	49.2	14.1	14775	24	AB199535	Mouse ischaemic co
20	48.4	13.9	2623	22	AAV17484	Mouse glucokinase
21	47.8	13.7	5318	21	AAV77094	Human ORFX ORF2649
22	46.8	13.4	35828	21	AAA29063	Murine TGF-beta b1
23	46.8	13.4	90050	21	AAZ91925	Wild type (C57BL/6
24	46.2	13.2	1459	22	AAV55245	Nucleotide sequenc
25	45.2	13.0	6789	22	AAV63436	Murine CD39-L4 gen
26	45.2	13.0	8212	24	AB199884	Mouse ischaemic co
27	45.2	13.0	13499	24	AAZ22571	Mouse FDRG (fibrin
28	44.6	12.8	48974	20	AAV55300	Mouse Presenilin-1
29	44.2	12.7	60	22	AAV81610	OST7 clone fragmen
30	44.2	12.7	60	22	AAV81611	Murine 45S pre rRN
31	43.6	12.5	4048	22	AAH24196	Mouse ageing inhibi
32	43.6	12.5	49999	20	AAZ23891	Murine LOBO genomi
33	43.6	12.5	49999	20	AAZ23896	Murine LOBO genomi
34	43	12.3	433	22	ABA77130	Proliferative glom
35	42.8	12.3	13146	18	AAV96719	Murine RENT1 genom
36	42.6	12.2	1104	20	AAZ10360	Partial genomic se
37	42.6	12.2	4612	24	AB199462	Mouse ischaemic co
38	42.4	12.1	6480	19	AAV99572	Mouse Friend virus
39	42	12.0	4580	17	AAV32034	Proliferation-inhi
40	42	12.0	90050	21	AAZ91925	Wild type (C57BL/6
41	41.4	11.9	8114	24	AAV15581	pPL5 fragment cont
42	41.4	11.9	11689	21	AAV87705	Mouse plakophilin-
43	41.2	11.8	467	23	AAV74838	DNA encoding novel
44	41.2	11.8	1445	22	AAV82696	Murine variant Zai
45	40.6	11.6	37339	22	AAV15612	Mouse osteocalcin

ALIGNMENTS

RESULT 1					
AAV32579	AAV32579	standard; cDNA; 1209 BP.			
XX	AAV32579;				
AC					
XX	23-SEP-1998	(first entry)			
DT					
XX					
DE	Human high voltage-dependent anion channel cDNA.				
XX					
KW	HACH; human high voltage-dependent anion channel; genomic mapping;				
KW	drug screening; proliferation disease; rheumatoid arthritis; tumour;				
KW	immuno-diagnosis; hypothalamus cDNA library; ss.				
XX					
OS	Homo sapiens.				
XX					
FH	Key	Location/Qualifiers			
FT	CDS	94..945			
FT		/*tag= a			
FT		/product= HACH			
XX					
PN	US5780235-A.				
XX					
PD	14-JUL-1998.				
XX					
PF	04-OCT-1996;	96US-0726227.			
XX					
PR	04-OCT-1996;	96US-0726227.			
XX					
PA	(INCY-) INCYTE PHARM INC.				
XX					
PI	Bandman O, Hillman JL;				

XX WPI: 1998-413045/35.
DR P-PSDB; AAW48908.
XX
PT New high voltage dependent anion channel protein and related nucleic
PT acid - vectors and transformed cells, useful for diagnosis and
PT treatment of tumours and other proliferative diseases
XX
PS Claim 3; Fig 1A-1B; 25pp; English.
XX
CC The present sequence represents the Human high voltage-dependent anion
CC channel (HACH) cDNA isolated from a hypothalamus cDNA library. Cells
CC transformed with HACH cDNA can be used to produce recombinant HACH
CC protein. HACH cDNA or its fragments, are claimed to be useful for
CC detecting and/or quantifying HACH gene expression (for diagnosis or
CC monitoring), as probes and primers for detecting genomic sequences
CC encoding HACH or related proteins. They are also claimed to be useful
CC in drug screening and genomic mapping. HACH protein or its activity
CC is claimed to be useful for inhibiting growth of tumours and for
CC treating other cell proliferation diseases, e.g. rheumatoid arthritis.
CC HACH protein and its fragments are also claimed to be useful for
CC screening binding agents for the protein, potential therapeutic
CC agents, and to raise antibodies. Antibodies can be useful for
CC diagnosing or monitoring HACH-related disorders, also therapeutically,
CC in competitive drug screens, and for affinity purification of the
CC HACH protein from natural sources.
XX
SQ Sequence 1209 BP; 342 A; 233 C; 301 G; 332 T; 1 other;

Query Match 37.4%; Score 130.6; DB 19; Length 1209;
Best Local Similarity 72.6%; Pred. No. 1.4e-34;
Matches 191; Conservative 0; Mismatches 68; Indels 4; Gaps 3;

QY 88 ATGGGCTGCNACTATGGGCTCACCTTCACCCANANGNGAGTAGCNGACGGTACTCTTGGC 147
DB 280 AAGGTCTGTAACATATGACTTACCTTCACCCAGAAATGGACACAGACAATACTCTAGGG 339
QY 148 ACAGACCTTTTGTGNGAATNTGCATGCTGANGGGTTNAACTGACTCTGATACAT 207
DB 340 ACAGAAATCTCTTGGGAGATAAG-TTGGCTGAAGGGTTGAACCTGACTCTGATACCAT 398
QY 208 ATTTNTACCATNCTCCNATCCCTTTAGTGCCATTTTCCCGCCTCTATTGCCNGNAT 267
DB 399 ATTTGTACCGAACACAGGAA--AGAGAGGTGGGAAATGGAAGGCTCCTATAAACGGGAT 456
QY 268 TGTNTNANTCTCGGACGATGATGTTGATNTNNATTTTCTGACCGACCATCTATGGCT-G 326
DB 457 TGTTTAGTGTGGCAGTATGTTGATATAGATTTTCTGACCAACCATCTATGGCTGG 516
QY 327 TCTGNNTGGCCTTTGAAGGTG 349
DB 517 GCTGTGTGGCCTTCGAAGGCTG 539

RESULT 2
AAT94469
ID AAT94469 standard; cDNA; 1393 BP.
XX
AC AAT94469;
XX
DT 03-MAR-1998 (first entry)
XX
DE Human Fchd545 gene differentially regulated in endothelial cells.
XX
KW Fchd545 gene; differential expression; endothelial cell; human;
KW shear stress; cardiovascular disease; atherosclerosis; ischaemia;
KW reperfusion; hypertension; restenosis; arterial inflammation;
KW therapy; diagnosis; drug screening; ss.
XX
OS Homo sapiens.
XX
FH Key Location/Qualifiers
FT CDS 90..941

FT
XX
XX WPI: 1997-424966/39.
PN P-PSDB; AAW36004.
XX
XX 21-AUG-1997.
PD
XX
XX 14-FEB-1997; 97WO-US02291.
PF
XX
PR 13-FEB-1997; 97US-0799910.
PR 16-FEB-1996; 96US-0011787.
XX
PA (MILL-) MILLENNIUM PHARM INC.
XX
PI Falb DA:
XX
DR WPI: 1997-424966/39.
DR P-PSDB; AAW36004.
XX

PT New genes differentially expressed in cardiovascular disease - used
PT for diagnosis, drug screening and treatment of cardiovascular
PT disease, e.g. atherosclerosis, restenosis, hypertension, etc
XX
PS Claim 1; Fig 3; 163pp; English.

CC Fchd545 is a novel human gene that is down-regulated in endothelial
CC cells subjected to laminar shear stress. Differential display was
CC used to detect genes that are differentially expressed in
CC endothelial cells subjected to fluid shear stress in vitro. Shear
CC stress is thought to be responsible for the prevalence of
CC atherosclerotic lesions in areas of unusual circulatory flow. Also
CC identified were novel gene fchd531 (see AAT94467), which is
CC up-regulated in endothelial cells under laminar and turbulent shear
CC stress, and fchd540 (see AAT94468), which is up-regulated in
CC endothelial cells under laminar shear stress. These genes provide a
CC fingerprint for the study of cardiovascular diseases, including
CC atherosclerosis, ischaemia/reperfusion, hypertension and restenosis.
CC Methods are provided for the diagnosis, monitoring in clinical
CC trials, screening for therapeutically effective compounds, and
CC treatment of cardiovascular diseases based on discoveries regarding
CC the expression patterns of novel genes fchd531, fchd540, fchd545,
CC fchd602 (see AAT94470) and fchd605 (see AAT96671). The fchd545 gene,
CC deposited as ATCC 69974, encodes a transmembrane protein (see
CC AAW36004) that provides an excellent target for detection of
CC cardiovascular disease states in diagnostic systems. The fchd545
CC gene is also expressed in the human heart, smooth muscle and
CC testis.
XX
SQ Sequence 1393 BP; 386 A; 269 C; 334 G; 404 T; 0 other;

Query Match 37.4%; Score 130.6; DB 18; Length 1393;
Best Local Similarity 72.6%; Pred. No. 1.4e-34;
Matches 191; Conservative 0; Mismatches 68; Indels 4; Gaps 3;

QY 88 ATGGGCTGCNACTATGGGCTCACCTTCACCCANANGNGAGTAGCNGACGGTACTCTTGGG 147
DB 276 AAGGTCTGTAACATATGACTTACCTTCACCCAGAAATGGACACAGACAATACTCTAGGG 335
QY 148 ACAGACCTTTTGTGNGAATNTGCATGCTGANGGGTTNAACTGACTCTGATACCAT 207
DB 336 ACAGAAATCTCTTGGGAGATAAG-TTGGCTGAAGGTTGAAGCTGACTCTGATACCAT 394
QY 208 ATTTNTACCATNCTCCNATCCCTTTAGTGCCATTTTCCCGCCTCTATTGCCNGNAT 267
DB 395 ATTTGTACCGAACACAGGAA--AGAGAGGTGGGAAATGGAAGGCTCCTATAAACGGGAT 452
QY 268 TGTNTNANTCTCGGACGATGATGTTGATNTNNATTTTCTGACCGACCATCTATGGCT-G 326
DB 453 TGTTTAGTGTGGCAGTATGTTGATATAGATTTTCTGACCAACCATCTATGGCTGG 512
QY 327 TCTGNNTGGCCTTTGAAGGTG 349
DB 513 GCTGTGTGGCCTTCGAAGGCTG 535

```
RESULT 3
AAZ50709
ID AAZ50709 standard; DNA; 1393 BP.
XX
AC AAZ50709;
XX
DT 31-MAY-2000 (first entry)
XX
DE Nucleotide sequence of human fchd545 gene.
XX
KW fchd545 gene; human; cardiovascular disease; oncogenic disorder;
KW diabetic retinopathy; fibroproliferative disorder; arteriosclerosis;
KW TGF-beta signalling pathway; TGF; Transforming growth factor;
KW pancreatic cancer; angiogenesis; inflammation; fibrosis; tumour growth;
KW vascularisation; cytostatic; antidiabetic; ophthalmological; ds.
XX
OS Homo sapiens.
XX
FH Key Location/Qualifiers
FT CDS 90..941
FT /*tag= a
FT /product= "fchd545 protein"
XX
PN WO200006206-A1.
XX
PD 10-FEB-2000.
XX
PF 30-JUL-1999; 99WO-US17394.
XX
PR 30-JUL-1998; 98US-0126640.
XX
PA (MILL-) MILLENNIUM PHARM INC.
XX
PI Falb DA;
XX
DR WPI: 2000-205414/18.
DR P-PSDB; AAY45015.
XX
PT Identifying substances for ameliorating symptoms of fibroproliferative
PT diseases or oncogenic related disorders -
XX
PS Examples; Fig 3; 214pp; English.
XX
CC The patent discloses methods for the treatment and diagnosis of
CC cardiovascular diseases by novel human genes which are differentially
CC expressed in different cardiovascular disease states. Compositions which
CC can modify TGF-beta signalling pathway are identified by screening.
CC These are used therapeutically to treat fibroproliferative and oncogenic
CC disorders, especially TGF (Transforming growth factor)-beta related
CC disorders, including diabetic retinopathy, arteriosclerosis, pancreatic
CC cancer, angiogenesis, inflammation, fibrosis, tumour growth and
CC vascularisation. The present sequence is fchd545 gene which is down
CC -regulated in endothelial cells subjected to shear stress can be used to
CC design cardiovascular disease treatment strategies. Depending on whether
CC the down-regulation has a pathogenic or protective effect treatment
CC methods can be designed to increase or decrease the activity of the
CC protein product of the gene.
XX
SQ Sequence 1393 BP; 406 A; 269 C; 333 G; 385 T; 0 other;

Query Match 37.4%; Score 130.6; DB 21; Length 1393;
Best Local Similarity 72.6%; Pred. No. 1.4e-34;
Matches 191; Conservative 0; Mismatches 68; Indels 4; Gaps 3;
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QY 208 ATTTACCATNCGNCTCCNATCCTTTAGTGCATTTTCCGGCCCTCTATGCGNGNAT 267
DB 395 ATTTGATCCGACACAGGAA--AGAAGAGTGGGAAATTTGAAGGCCCTCTATAACGGGAT 452
QY 268 TGTNNANNTCTCGGCAGTAATGTGATNTNNAATTTTCTGACCGGACCATCTATGGCT-G 326
DB 453 TGTTTAGTGTGGCAGTAATGTGATATAGATTTTCTGACCAACCATCTATGGCTGG 512
QY 327 TCTGNNTTGGCCTTTGAAGGTTG 349
DB 513 GCTGTGTTCGCCCTTCGAAGGGTG 535

RESULT 4
AAZ57719
ID AAZ57719 standard; CDNA; 1414 BP.
XX
AC AAZ57719;
XX
DT 16-JUL-1999 (first entry)
XX
DE Voltage-dependent anion channel CBMAAD07 coding sequence.
XX
KW Human; voltage-dependent anion channel; CBMAAD07; antibody; antagonist;
KW cancer; spontaneous abortion; infertility; ss.
XX
OS Homo sapiens.
XX
PN WO9921990-A1.
XX
PD 06-MAY-1999.
XX
PF 29-OCT-1997; 97WO-CN00118.
XX
PR 29-OCT-1997; 97WO-CN00118.
XX
PA (UYSH-) UNIV SHANGHAI SECOND MEDICAL.
XX
PI Wang Y, Zhang Q;
XX
DR WPI: 1999-303016/25.
DR P-PSDB; AAY07222.
XX
PT CBMAAD07, a human voltage-dependent anion channel protein, useful in
PT the treatment and diagnosis of microsomal and neurological disorders
XX
PS Claim 4; Page 7-8; 31pp; English.
XX
CC This sequence represent the coding sequence for a novel human
CC voltage-dependent anion channel designated CBMAAD07. The protein,
CC antibodies and (ant)agonists to it can be used for treating, e.g.
CC cancer, spontaneous abortion and infertility.
XX
SQ Sequence 1414 BP; 419 A; 270 C; 334 G; 391 T; 0 other;

Query Match 37.4%; Score 130.6; DB 20; Length 1414;
Best Local Similarity 72.6%; Pred. No. 1.5e-34;
Matches 191; Conservative 0; Mismatches 68; Indels 4; Gaps 3;
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QY 88 ATGGGCTGCNACTATGGGCTCACCCTTCACCCANANGNGAGTACNGAGGTACTCTTGGG 147
DB 276 AAGGTCTGTAATGAGACTTACCTTCACCCAGAAATGGAACACAGACATACTCTAGGG 335
QY 148 ACAGACCTTTTGTGNGAGAAATNTGCATGGCTGANGGCTTNAACCTGACTCTGCATACCAT 207
DB 336 ACAGAAATCTCTGGGAGATAAG-TTGGCTGAAGGGTTGAACACTGACTCTGTATACCAT 394
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QY 88 ATGGGCTGCNACTATGGGCTCACCCTTCACCCANANGNGAGTACNGAGGTACTCTTGGG 147
DB 286 AAGGTCTGTAATGAGACTTACCTTCACCCAGAAATGGAACACAGACATACTCTAGGG 345
QY 148 ACAGACCTTTTGTGNGAGAAATNTGCATGGCTGANGGCTTNAACCTGACTCTGCATACCAT 207
DB 346 ACAGAAATCTCTGGGAGATAAG-TTGGCTGAAGGGTTGAACACTGACTCTGTATACCAT 404
QY 208 ATTTNACCATNCGNCTCCNATCCTTTAGTGCCATTTTCCGGCCCTCTATGCGNGNAT 267
DB 405 ATTTGATCCGACACAGGAA--AGAAGAGTGGGAAATTTGAAGGCCCTCTATTAACGGGAT 462
QY 268 TGTNNANNTCTCGGCAGTAATGTGATNTNNAATTTTCTGACCGGACCATCTATGGCT-G 326
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CC responsible for genetic disorders or other traits to assess biodiversity
CC and to produce other types of data and products dependent on DNA and
CC amino acid sequences. AAS64197-AAS94564 represent novel human
CC diagnostic coding sequences of the invention.
CC Note: The sequence data for this patent did not appear in the printed
CC specification, but was obtained in electronic format directly from WIPO
CC at ftp.wipo.int/pub/published_pcl_sequences.

XX Sequence 1672 BP; 423 A; 376 C; 451 G; 422 T; 0 other;

Query Match 30.1%; Score 105; DB 23; Length 1672;
Best Local Similarity 64.0%; Pred. No. 1e-25;
Matches 181; Conservative 0; Mismatches 98; Indels 4; Gaps 3;

OY 66 GCGNCGCTANAGNCCNCAATATCATGGGCTGCNACTATGGGCTCACCTTCACCCANANGNG 125
Db 534 GCGAACCCTAGAACAACCAATATAAGGCTCTGTACTATGGACTTACCCCTCACCAGAAATG 593
OY 126 GAGTACNGACGGTACTCT-TGGGACAGACCTTTTGTGNGAATAINTGCATGGCTGANGGG 184
Db 594 GAACACAGACAATACTCTTAAGGACAGACAAGAAATCTTTGGAGAAATAAGTTGGCTGAAGGG 653
OY 185 TTNAAACTGACTCTCGATACCAATATTNTACCATNCNCTCCNATCCTTTAGTGCCATTT 244
Db 654 TTGAAACTGACTCTTGATATACCAATATTGTGACCAGACAGAGAA--AGAAGAGTGGAAT 711
OY 245 TCCCGGCTCTCTATTTGCCNGNATGNTNANNTCTCGGCAGTAATGTGATNTNATTTT 304
Db 712 TGAAGGCTCTCTATTAACGGAATGTTTAGTGTGGCAGTAATGTTGATATAGATTTT 771
OY 305 CTGGACCGACCATCTATGGCT-GTCTGNNTTGGCCTTTGAAGG 346
Db 772 CTGGACCAACCATCTATGGCTGGCTGTGCTTGGCCTTCGGAAG 814

RESULT 7
AAC01263

ID AAC01263 standard; cDNA; 435 BP.

XX AAC01263;

DT 06-OCT-2000 (first entry)

XX Human secreted protein 5' EST, SEQ ID NO: 1261.

DE Human secreted protein 5' EST, SEQ ID NO: 1261.
XX
KW Human; 5' EST; expressed sequence tag; secreted protein; cDNA isolation;
KW gene therapy; chromosome mapping; ss.

XX Homo sapiens.

OS
PN EP1033401-A2.

XX
PD 06-SEP-2000.

XX
PF 21-FEB-2000; 2000EP-0200610.

XX
PR 26-FEB-1999; 99US-0122487.

XX
PA (GEST) GENSET.

XX
PI Dumas Milne Edwards J, Duclert A, Giordano J;

XX
DR WPI; 2000-500381/45.

DR P-PSDB; AAC01257.

XX
PT New nucleic acid that is a 5' expressed sequence tag (5' EST) for
PT obtaining cDNAs and genomic DNAs that correspond to 5'ESTs and for
PT diagnostic, forensic, gene therapy and chromosome mapping procedures -

XX
PS Claim 1; SEQ ID 1261; 71pp + CD-ROM; English.

XX
CC The present sequence is one of a large number of 5' ESTs derived from
CC mRNAs encoding secreted proteins. An ORF has been identified within the

CC sequence. The 5' ESTs were prepared from total human RNAs or polyA+ RNAs
CC derived from 30 different tissues. EST sequences usually correspond
CC mainly to the 3' untranslated region (UTR) of the mRNA because they are
CC often obtained from oligo-dT primed cDNA libraries. Such ESTs are not
CC well suited for isolating cDNA sequences derived from the 5' ends of
CC mRNAs and even in those cases where longer cDNA sequences have been
CC obtained, the full 5' UTR is rarely included. 5' ESTs are derived from
CC mRNAs with intact 5' ends and can therefore be used to obtain full length
CC cDNAs and genomic DNAs. 5' ESTs are also used in diagnostic, forensic,
CC gene therapy and chromosome mapping procedures. They are used to obtain
CC upstream regulatory sequences and to design expression and secretion
CC vectors.

XX Sequence 435 BP; 127 A; 83 C; 114 G; 108 T; 3 other;

Query Match 22.3%; Score 77.8; DB 21; Length 435;
Best Local Similarity 76.7%; Pred. No. 1.3e-16;
Matches 99; Conservative 1; Mismatches 28; Indels 1; Gaps 1;

OY 88 ATGGCTGCNACTATGGGCTCACTTCACCCANANGNGAGTACNGACGTACTCTTGGG 147
Db 308 AAGGCTGTAACTATGACTTACTTCACTTCACCCAGAAATGGAACACAGAYAAATACTTACGG 367
OY 148 ACAGACCTTTTGTGNGAATNTGCATGGCTGANGGGTTNAACCTGACTCTGCATACCAT 207
Db 368 ACAGAAATCTCTTGGGAGAAATAG-TTGGCTGAAGGGTTGAACCTGACTCTTGATACCAT 426
OY 208 ATTTNTACC 216
Db 427 ATTTGTACC 435

RESULT 8
AAS74836

ID AAS74836 standard; cDNA; 473 BP.

XX AAS74836;

DT 13-FEB-2002 (first entry)

XX DNA encoding novel human diagnostic protein #10640.

DE Human; chromosome mapping; gene mapping; gene therapy; forensic;
KW food supplement; medical imaging; diagnostic; genetic disorder; ss.

XX Homo sapiens.

OS
PN WO200175067-A2.

XX
PD 11-OCT-2001.

XX
PF 30-MAR-2001; 2001WO-US08631.

XX
PR 31-MAR-2000; 2000US-0540217.

XX
PR 23-AUG-2000; 2000US-0649167.

XX
PA (HYSE-) HYSEQ INC.

XX
PI Drmanac RT, Liu C, Tang YT;

XX
DR WPI; 2001-639362/73.

DR P-PSDB; ABG10649.

XX
PT New isolated polynucleotide and encoded polypeptides, useful in
PT diagnostics, forensics, gene mapping, identification of mutations
PT responsible for genetic disorders or other traits and to assess
PT biodiversity -

XX
PS Claim 1; SEQ ID No 10640; 103pp; English.

XX
CC The invention relates to isolated polynucleotide (I) and
CC polypeptide (II) sequences. (I) is useful as hybridisation probes,
CC polymerase chain reaction (PCR) primers, oligomers, and for chromosome

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XX		WO9412162-A.
XX		
XX		09-JUN-1994.
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XX		01-DEC-1993; 93WO-F100514.
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XX		01-DEC-1992; 92US-0988427.
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XX		(ALAN/) ALANIN-KURKI L M.
XX		(AUVI/) AUVINEN P O V.
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XX		Alanen-Kurki LM, Auvinen POV, Jaakkola PM, Jalkanen MT;
XX		Leppaesm, Mali MS, Vihinen TA, Waerri AM;
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XX		WPI, 1994-199926/24.
XX		P-PSDB; AAR55276.
XX		
XX		Syndecan stimulation of cellular differentiation - useful for
XX		decreasing tumour growth used to promote hair growth
XX		
XX		Disclosure; Page 22-39; 65pp; English.
XX		
XX		The mouse syndecan gene enhancer, located 8-10 kb upstream from the
XX		initiation site, is given in AAQ67901. Manipulation of the enhancer
XX		can be used either to slow or prevent tumor growth or to promote
XX		differentiation of specific cell types, e.g. epidermal cells to
XX		promote hair formation. The complete mouse syndecan gene and its
XX		encoded protein are given in AAQ67902 and AAR55276.
XX		
XX		Sequence 26700 BP; 5742 A; 6559 C; 7233 G; 7165 T; 1 other;
XX		
XX		Query Match 14.6%; Score 51; DB 15; Length 26700;
XX		Best Local Similarity 66.0%; Pred. No. 1.6e-06;
XX		Matches 66; Conservative 0; Mismatches 34; Indels 0; Gaps 0;
XX		
QY	1 TATTATATGTAAGTACNCTGTAGCAGTTGTNNGACACACTCTACGAGGCGNCAGATCTC 60	
Db	1519 TATTATATGTAAGTACACTGTAGCTGTCTTCAGACACTCCAGAGAGCGCCAGATCTC 1578	
QY	61 ATTGTGGGNGGCTANAGNCNCATATCATGTGGCTGCNACT 100	
Db	1579 GTTATGATGTTGTTGTGAGCACCATGTGTTGCTGGGAATT 1618	
XX		
XX		RESULT 11
XX		AAV15946
XX		ID AAV15946 standard; DNA; 26700 BP.
XX		AC AAV15946;
XX		
XX		28-MAY-1998 (first entry)
XX		
XX		Mouse syndecan gene sequence.
XX		
XX		Syndecan; tumour suppression; tissue regeneration; enhancement;
XX		mouse; wound healing; ds.
XX		

OS	Mus sp.	Location/Qualifiers
XX	Key	4378..24421
FH	CDS	
FT		/tag= a
FT		/product= "syndecan protein"
FT		/note= "contains introns"
FT		1..4377
FT	intron	/tag= b
FT		/number= 1
FT	exon	4378..4443
FT		/tag= c
FT		/number= 1
FT	intron	4444..22025
FT		/tag= d
FT		/number= 2
FT	exon	22026..22106
FT		/tag= e
FT		/number= 2
FT	intron	22107..23000
FT		/tag= f
FT		/number= 3
FT	exon	23001..23483
FT		/tag= g
FT		/number= 3
FT	intron	23484..23904
FT		/tag= h
FT		/number= 4
FT	exon	23905..24039
FT		/tag= i
FT		/number= 4
FT	intron	24040..24250
FT		/tag= j
FT		/number= 5
FT	exon	24251..24418
FT		/tag= k
FT		/number= 5
FT	intron	24422..26700
FT		/tag= l
FT		/number= 6
XX		
PN	US5726058-A.	
XX		
PD	10-MAR-1998.	
XX		
PF	07-JUN-1995;	95US-0472217.
XX		
PR	07-MAR-1994;	94US-0206186.
PR	01-DEC-1992;	92US-0988427.
PR	01-DEC-1993;	93WO-F100514.
PR	07-JUN-1995;	95US-0472217.
XX		
PA	(ALAN/) ALANEN-KURKI L.	
PA	(AUVI/) AUVINEN P.	
PA	(JAAK/) JAAKKOLA P.	
PA	(JALK/) JALKANEN M.	
PA	(LEPP/) LEPPAE S.	
PA	(MALI/) MALI M.	
PA	(VIHI/) VIHINEN T.	
PA	(WAER/) WAERRI A.	
XX		
XX	Alanen-kurki L, Auvinen P, Jaakkola P, Jalkanen M;	
PI	Leppae S, Mali M, Vihinen T, Waerri A;	
XX		
DR	WPI; 1998-192770/17.	
XX		
DR	P-PSDB; AAW47156.	
XX		
PT	New mouse syndecan gene sequences - useful for, e.g. suppressing	
PT	tumour growth or promoting tissue regeneration in processes such as	
PT	wound healing	
XX		
PS	Claim 2; Fig 2A-O; 48pp; English.	
XX		

CC This is the mouse syndecan gene sequence. A 350 base pair fragment
CC (AAV15948) of a purified 2196 base pair DNA molecule (AAV15947) enhances
CC the expression of a gene operably linked to the promoter of the mouse
CC syndecan gene in 3T3 cells following treatment with TGF- beta and bFGF
CC when the fragment is operably linked to the promoter. A purified DNA
CC molecule comprising a portion of the nucleotide residues 3538-3888 of
CC the mouse syndecan genomic sequence suppresses expression of a gene
CC operably linked to the promoter of the mouse syndecan gene in 5115 cells
CC treated with testosterone. Host cells can be transfected with vectors
CC which contain either the enhancing or suppressing DNA molecules. The
CC products may be used to alter the differentiated state of a host cell by
CC altering its expression of syndecan, e.g. to induce and regulate
CC syndecan expression, especially in cells which exhibit a malignant
CC phenotype, regardless of the origin of transformation. The products can
CC be used to produce therapeutics for suppressing tumour growth. They may
CC enhance the syndecan expression in a host cell, by enhancing its gene
CC transcription, especially in malignant or normal cells, and therefore
CC promote tissue regeneration, especially in processes such as wound
CC healing.

CC XX Sequence 26700 BP; 5742 A; 6559 C; 7233 G; 7165 T; 1 other;

Query Match 14.6%; Score 51; DB 19; Length 26700;
Best Local Similarity 66.0%; Pred. No. 1.6e-06;
Matches 66; Conservative 0; Mismatches 34; Indels 0; Gaps 0;

OY 1 TATTATATGTAAGTACNCTGTAGACAGTGTGNNGACACACTCTACGAGGGCNCAGATCTC 60
DB 1519 TATTATATGTAAGTACACTGTAGCTGTCTTCAGACACTCCAGAGAGGGCGCAGATCTC 1578
OY 61 ATTGTGGGNGGCTANAGNCCNCATATCATGGGCTGCNACT 100
DB 1579 GTTATGATGCTGTCTGAGCACCACCATGCTGCTTGGGAATTT 1618

RESULT 12
AAV81283
ID AAV81283 standard; DNA; 26700 BP.
XX AC AAV81283;
XX 11-MAR-1999 (first entry)
XX Mouse syndecan-1 encoding DNA.
XX Syndecan-1; tumour; ectodomain; epithelial; mesenchymal; breast; mouse;
KW endometrial tumour; prostatic tumour; oestrogenic; androgenic; steroid;
KW glioma; myeloma; carcinoma; sarcoma; lymphoma; adenoma; ss.
OS Mus sp.

XX FH Key Location/Qualifiers
FT CDS 4378..24421
FT /*tag= a
FT /product= "mouse syndecan-1"
FT /note= "contains introns"
FT 4378..4443
FT /*tag= b
FT /number= 1
FT 4444..22025
FT /*tag= c
FT /number= 1
FT 22026..22107
FT /*tag= d
FT /number= 2
FT 22108..23001
FT /*tag= e
FT /number= 2
FT 23002..23483
FT /*tag= f
FT /number= 3
FT 23484..23904
FT /*tag= g
FT Intron

FT /number= 3
FT exon 23905..24040
FT /*tag= h
FT /number= 4
FT Intron 24041..24251
FT /*tag= i
FT /number= 4
FT exon 24252..2418
FT /*tag= j
FT /number= 5

PN US5851993-A.
XX 22-DEC-1998.
PD 07-JUN-1995; 95US-0488199.
XX 07-JUN-1995; 95US-0488199.
PR 07-JUN-1995; 95US-0488199.
PR 13-JUN-1994; 94US-0258862.
XX (BIOT-) BIOTIE THERAPIES LTD.
XX Jalakanen M, Mali M;
XX WPI; 1999-104635/09.
DR P-PSDB; AAW95198.

XX Reducing growth of tumour cells - with ectodomain of syndecan
PT applied to the extracellular environment to induce a more
PT differentiated phenotype, particularly for hormone-dependent breast,
PT endometrial or prostatic cancers
XX Disclosure; Fig 2A-M; 48pp; English.

XX The invention relates to a method of reducing the growth of tumour cells
CC which comprises supplying a human syndecan ectodomain to the environment
CC around the cells causing them to develop a more differentiated phenotype.
CC The method is used to suppress tumour cells of epithelial, mesenchymal,
CC pre-B or plasma cell origin, especially breast endometrial or prostatic
CC tumours, and particularly those responsive to oestrogenic or androgenic
CC steroid. More generally it can be used to treat malignant or non-
CC malignant tumours, particularly those characterised by loss of syndecan,
CC e.g. gliomas, myelomas, carcinomas, sarcomas, lymphomas and adenomas. The
CC present sequence represents the DNA sequence encoding a mouse syndecan-1.
XX Sequence 26700 BP; 5742 A; 6559 C; 7233 G; 7165 T; 1 other;

Query Match 14.6%; Score 51; DB 20; Length 26700;
Best Local Similarity 66.0%; Pred. No. 1.6e-06;
Matches 66; Conservative 0; Mismatches 34; Indels 0; Gaps 0;

OY 1 TATTATATGTAAGTACNCTGTAGACAGTGTGNNGACACACTCTACGAGGGCNCAGATCTC 60
DB 1519 TATTATATGTAAGTACACTGTAGCTGTCTTCAGACACTCCAGAGAGGGCGCAGATCTC 1578
OY 61 ATTGTGGGNGGCTANAGNCCNCATATCATGGGCTGCNACT 100
DB 1579 GTTATGATGCTGTCTGAGCACCACCATGCTGCTTGGGAATTT 1618

RESULT 13
AAV85695/c
ID AAV85695 standard; cDNA; 178 BP.
XX AC AAV85695;
XX 06-SEP-1999 (first entry)
XX Novel cDNA sequence from a mouse blastocyst cDNA library.
DE Mouse; blastocyst; cDNA library; ss.
XX Mus sp.

XX JP11164691-A.
PN 22-JUN-1999.
XX
XX 14-APR-1998; 98JP-0103115.
PF 03-OCT-1997; 97JP-0271781.
XX
XX (RIKA) RIKAGAKU KENKYUSHO.
XX WPI; 1999-411831/35.
DR
XX
XX New blastocyst cDNA - useful for library construction
PT
XX
XX Claim 3; Page 26; 41pp; Japanese.
PS
XX
XX AAX85621-X85746 represent novel cDNA sequences that are isolated from a
CC mouse blastocyst cDNA library. The cDNA library was constructed from
CC C57Bl/6 mice. The sequence can be used as a source of primers, probes
CC and complementary DNA sequences.
XX
SQ Sequence 178 BP; 55 A; 42 C; 39 G; 42 T; 0 other;

Query Match 14.3%; Score 50; D 20; Length 179;
Best Local Similarity 72.0%; Pred. No. 3..e-07;
Matches 59; Conservative 0; Mismatches 23; Indels 0; Gaps 0;

QY 1 TATATATGTAGTACNCTGTAGCAGTTGTNNGACACACTCTACGAGGGCNCAGATCTC 60
Db 160 TATATATGTAGTACACTGTAGCTGTCTTCAGACACTCCAGAGAGGGCATCAGATCTC 101
QY 61 ATTGTGGGNGGCTANAGNCNC 82
Db 100 ATTACGGGTGGTGTGAGCCAC 79

RESULT 14
AAV84328/c
ID AAV84328 standard; DNA; 5889 BP.

AC AAV84328;

DT 26-APR-1999 (first entry)

DE Mouse A-myb genomic DNA.

KW A-myb gene; oncogene; transgenic animal; null mutant; stem cell;
KW spermatogenesis; infertility; knockout mouse; animal model; ss.

OS Mus sp.

PN WO9846726-A1.

PD 22-OCT-1998.

PF 07-APR-1998; 98WO-US06896.

PR 15-APR-1997; 97US-0043353.

PA (UTEM) UNIV TEMPLE.

PI Halton K, Reddy EP, Toscani A;

DR WPI; 1999-080737/07.

PT Transgenic non-human animal with disrupted A-myb locus in the genome
PT - useful as model for male infertility and for studying
PT spermatogenesis

PS Example 1; Page 51-55; 83pp; English.

CC This is the nucleotide sequence of a genomic clone of the mouse

CC A-myb gene. The gene was isolated by screening a lambda DASH mouse
CC genomic library derived from the 129/J mouse strain, using a probe
CC derived from the 5' end of an A-myb cDNA clone that encodes the
CC DNA binding domain of the protein. Positive clones that contained
CC the 5.9 kb HindIII fragment were subcloned into pGEM 7zf(+) plasmid
CC vector. The 5.9 kb fragment contains exons 3, 4 and 5 of the gene
CC that code for the 5' end of the DNA binding domain of the protein.
CC The genomic clone was deposited as NRRL B-21575. The invention
CC provides non-human animals in which expression of the A-myb gene is
CC suppressed. A transgenic non-human animal (especially a mouse), or
CC stem cell, having a disrupted A-myb gene in the genome, is claimed.
CC Also new are: (1) spermatogonia comprising recombinant DNA encoding
CC a functional A-myb polypeptide; (2) a targeting vector for
CC functionally disrupting an A-myb gene comprising a polynucleotide
CC where one part of the polynucleotide contains a sequence homologous
CC to sequences in or flanking an A-myb gene, and which, when
CC integrated into the corresponding A-myb locus, functionally
CC disrupts the A-myb gene; and (3) methods of restoring fertility to
CC a subject who is infertile due to a defect in the A-myb locus, by
CC administration of A-myb or DNA encoding A-myb. Male A-myb -/-
CC animals can be used as models for male infertility and for studying
CC spermatogenesis. They can be used to test 'rescue' constructs and
CC other agents to treat male infertility.

SQ Sequence 5889 BP; 1828 A; 922 C; 1089 G; 2050 T; 0 other;

Query Match 14.3%; Score 50; DB 20; Length 5889;
Best Local Similarity 72.0%; Pred. No. 1.7e-06;
Matches 59; Conservative 0; Mismatches 23; Indels 0; Gaps 0;

QY 1 TATATATGTAGTACNCTGTAGCAGTTGTNNGACACACTCTACGAGGGCNCAGATCTC 60
Db 2148 TATATATGTAGTACACTGTAGCTGTCTTCAGACACCCAGAGAGGGCATCAGATCTC 2089
QY 61 ATTGTGGGNGGCTANAGNCNC 82
Db 2088 ATTATGATGTTGTGAGCCAC 2067

RESULT 15
AAF63435
ID AAF63435 standard; DNA; 6094 BP.

AC AAF63435;

DT 14-MAY-2001 (first entry)

DE Murine CD39-L2 genomic DNA sequence.

KW Human CD39-like protein; apyrase; NDPase; platelet function inhibitor;
KW myocardial infarction; cerebral ischaemia; angina; arterial thrombosis;
KW cerebral artery thrombosis; platelet aggregation; inflammation;
KW apoptosis; autoimmune disorder; neurological disorder;
KW Alzheimer's disease; Parkinson's disease; cancer; CD39-L2; ds.

OS Mus sp.

PN WO200110205-A1.

PD 15-FEB-2001.

PF 09-AUG-2000; 2000WO-US21790.

PR 09-AUG-1999; 99US-0370265.

PR 11-JAN-2000; 2000US-0481238.

PR 25-APR-2000; 2000US-0557800.

PR 26-MAY-2000; 2000US-0583231.

PI 30-JUN-2000; 2000US-0608285.
(HYSE-) HYSEQ INC.
Ford J, Mulero JJ, Yeung G;

DR WP1: 2001-147489/15.

XX Polynucleotides encoding human CD39-like polypeptides, with apyrase
PT and/or NDPase activity, which are useful in the treatment of
PT pathological conditions caused by thrombosis (e.g. myocardial
PT infarction) and inflammatory disorders -

XX Example 21; Page 106-108; 203pp; English.

PS This invention relates to polynucleotides encoding human CD39-like
XX polypeptides with apyrase and/or NDPase activity. The polypeptides having
CC polypeptides with apyrase, activity are useful for inhibiting platelet
CC ATPase, including NDPase, be used in the prophylaxis or treatment of
CC function and can therefore be used by or involving thrombosis or excessive
CC pathological conditions caused by or involving thrombosis, such as myocardial
CC coagulation or excessive platelet aggregation, such as myocardial
CC infarction, cerebral ischaemia, angina, arterial thrombosis, cerebral
CC artery thrombosis or intracardiac thrombosis, and conditions associated
CC with venous thrombosis. CD39-L4 and CD39-L2 polypeptides are useful in
CC modulating disease states (including platelet aggregation, inflammation
CC and apoptosis) associated with ADP or other purinergic signalling by
CC reducing the levels of NDPs. The polypeptides are also useful for
CC prophylaxis or treatment of inflammation related disorders, such as
CC disorders involving sepsis or systemic inflammatory response syndrome or
CC SIRS (and associated conditions such as fever, tachycardia, tachypnea,
CC cytokine overstimulation); autoimmune disorders such as thrombosis,
CC atherosclerosis, acute pancreatitis, dermatitis, including psoriasis,
CC cirrhosis, reperfusion injury, asthma, multiple sclerosis, arthritis,
CC neurological disorders including neurodegenerative diseases, epilepsy,
CC depression, Alzheimer's disease, Parkinson's disease, Huntington's
CC disease, and amyotrophic lateral sclerosis; and cancer. The present
CC sequence represents the murine CD39 like protein CD39-L2 genomic DNA
CC sequence.

XX Sequence 6094 BP; 1589 A; 1471 C; 1445 G; 1504 T; 85 other;

SQ Query Match 14.3%; Score 50; DB 22; Length 6094;
Best Local Similarity 72.0%; Pred. No. 1.7e-06;
Matches 59; Conservative 0; Mismatches 23; Indels 0; Gaps 0;

QY 1 TATTATATGTAGTACNCTGTAGCAGTGTGTTNGACACACTCTACGAGGCGCCAGATCTC 60
Db 3329 TATTATATGTAGTACACTGTGCTCTTCACACACTCCAGAAGCGCATCAGATCTC 3388
QY 61 ATTGTGGGNGGCTANAGNCNC 82
Db 3389 ATTACGGGTGGTGTGAGCCAC 3410

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